Prefrontal Cortex Regulates Chronic Stress-Induced Cardiovascular Susceptibility

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Key Points

- Knockdown of glutamate release from infralimbic cortex increases heart rate and arterial pressure reactivity
- Reduced infralimbic glutamate release increases cumulative arterial pressure during chronic variable stress
- Decreased infralimbic glutamate output leads to vascular dysfunction after chronic stress
- These functional changes associate with histological indicators of cardiac hypertrophy, as well as vascular hypertrophy and fibrosis.

Running Title

Cortical regulation of cardiovascular responses to stress
Abstract

The medial prefrontal cortex (mPFC) is necessary for appropriate appraisal of stressful information, as well as coordinating visceral and behavioral processes. However, prolonged stress impairs mPFC function and prefrontal-dependent behaviors. Additionally, chronic stress induces sympathetic predominance, contributing to health detriments associated with autonomic imbalance. Previous studies identified a subregion of rodent prefrontal cortex, infralimbic cortex (IL), as a key regulator of neuroendocrine-autonomic integration after chronic stress, suggesting that IL output may prevent chronic stress-induced autonomic imbalance. In the current study, we tested the hypothesis that the IL regulates hemodynamic, vascular, and cardiac responses to chronic stress. To address this hypothesis, a viral-packaged siRNA construct was used to knockdown vesicular glutamate transporter 1 (vGluT1) and reduce glutamate packaging and release from IL projection neurons. Male rats were injected with a vGluT1 siRNA-expressing construct or GFP control into the IL and then remained as unstressed controls or were exposed to chronic variable stress (CVS). IL vGluT1 knockdown increased heart rate and mean arterial pressure (MAP) reactivity, while CVS increased chronic MAP only in siRNA-treated rats. In a separate cohort, CVS and vGluT1 knockdown interacted to impair both endothelial-dependent and endothelial-independent vasoreactivity ex vivo. Furthermore, vGluT1 knockdown and CVS increased histological markers of fibrosis and hypertrophy. Thus, knockdown of glutamate release from IL projection neurons indicates that these cells are necessary to prevent the enhanced sympathetic responses to stress that promote susceptibility to cardiovascular pathophysiology. These findings provide evidence for a neurobiological mechanism mediating the relationship between stress and poor cardiovascular health outcomes.

Keywords: infralimbic cortex, heart rate, blood pressure, vasoreactivity
Introduction

Stress, a real or perceived threat to homeostasis or well-being, elicits behavioral and physiological responses to promote organismal adaptation (de Kloet et al., 2005; Myers et al., 2014b). However, prolonged stress exposure has deleterious effects on health, increasing susceptibility to cardiovascular, psychiatric, and metabolic disorders (Grippo & Johnson, 2009; Binder & Nemeroff, 2010; Wardle et al., 2011; Sgoifo et al., 2015). In fact, chronic psychosocial stress predicts the incidence of cardiovascular disease, cardiac-related morbidity and mortality, and doubles the risk of myocardial infarction (Barefoot et al., 1996; Yusuf et al., 2004). Exaggerated heart rate (HR) reactivity to acute stress also predicts poor cardiovascular outcomes, including hypertension, ventricular hypertrophy, and atherosclerosis (Chida & Steptoe, 2010). Although the biological mechanisms mediating the relationship between stress and cardiovascular health are not completely understood, adverse outcomes likely result from prolonged exposure to neural and endocrine stress mediators.

The initial appraisal of psychological stressors largely occurs in the limbic system, a network of interconnected structures spanning the forebrain. The medial prefrontal cortex (mPFC) is a key limbic cortical structure mediating stress appraisal, emotion, and cognition (Damasio, 1996; Wood & Grafman, 2003; Myers-Schulz & Koenigs, 2012; McKlveen et al., 2015). Moreover, activity within a specific region of the ventral mPFC, the subgenual cingulate cortex (BA25), associates with sadness in healthy controls (Liotti et al., 2000), as well as pathological depression in treatment-resistant patients (Mayberg et al., 2005). Recent neuroimaging studies have also identified the ventral mPFC as a component of the central autonomic network that
responds to and coordinates visceral functions, including stress-evoked blood pressure reactivity (Gianaros & Sheu, 2009; Beissner et al., 2013; Gianaros & Wager, 2015; Shoemaker et al., 2015). The rodent homolog of BA25 is infralimbic cortex (IL) (Öngür et al., 2003; Uylings et al., 2003; Vertes, 2004). This subregion of ventral mPFC provides inputs to stress-integrative nuclei, including the posterior hypothalamus and brainstem autonomic centers (Gabbott et al., 2005; Myers et al., 2014a, 2016; Wood et al., 2019). Our previous studies reduced glutamate outflow from the IL in rats undergoing chronic variable stress (CVS), promoting hyperactivation of the hypothalamic-pituitary-adrenal axis (HPA) axis (Myers et al., 2017). As glutamate release from the IL is a key regulator of acute and chronic neuroendocrine reactivity, we hypothesized that IL output may prevent chronic stress-induced autonomic imbalance and associated cardiovascular susceptibility.

To address this hypothesis, a lentiviral-packaged small interfering RNA (siRNA) targeting vesicular glutamate transporter 1 (vGluT1) was injected in the IL. This approach selectively reduces vGluT1 expression in IL glutamate neurons, preventing the packaging and release of glutamate from presynaptic terminals (Wojcik et al., 2004; Schuske & Jorgensen, 2004; Myers et al., 2017). Animals were then exposed to CVS to examine interactions between chronic stress and hypo-functionality of ventral mPFC in terms of cardiovascular reactivity, arterial function, and remodeling of the vasculature and myocardium. These studies identified the necessity of the IL for reducing hemodynamic, vascular, and cardiac consequences of prolonged stress. Additionally, this work points toward a neurobiological mechanism mediating the relationship between stress and cardiovascular health.
Methods

Animals

Adult male Sprague-Dawley rats were obtained from Harlan (Indianapolis, IN) with weights ranging from 250-300 g. Rats were housed individually in shoebox cages in a temperature- and humidity-controlled room with a 12-hour light-dark cycle (lights on at 0600h, off at 1800h) and food and water ad libitum. All procedures and protocols were approved by Institutional Animal Care and Use Committees and comply with the National Institutes of Health Guidelines for the Care and Use of Laboratory Animals.

Experiment 1

Design

For experiment 1, 32 rats (n = 8/group) were injected with either a lentiviral-packaged construct coding for vGluT1 siRNA or GFP as a control. After instrumentation with radiotelemetry devices, half of the animals were exposed to 14 days of CVS with the rest of the animals remaining as No CVS controls. Home cage cardiovascular parameters were continuously monitored in all rats throughout the 14-day period of CVS. On the morning of day 15, all rats were subject to an acute novel restraint to examine hemodynamic stress reactivity.

Viral construct

A lentivirus transfer vector, based on a third-generation, self-inactivating transfer vector was constructed as previously described (Myers et al., 2017; Wood et al., 2019). Briefly, a 363-bp piece of DNA from the rat vGluT1 complementary DNA was
synthesized that included 151 bp of the 3’ coding region and 212 bp of the 3’ untranslated region. This corresponds to nucleotides 1656 to 2018 of GenBank accession no. NM_053859. This is a region of low homology with vGluT2 and vGluT3 and avoids all the putative transmembrane domains of the transporter. The fragment was cloned in antisense orientation into a lentivirus transfer vector that expressed an enhanced green fluorescent protein (GFP) reporter. This vector uses the phosphoglycerate kinase-1 promoter, which expresses well in rat brain and is primarily neuronal (Grillo et al., 2015; Myers et al., 2017). A control virus was constructed similarly, using a transfer vector with the phosphoglycerate kinase-1 promoter driving expression of enhanced GFP alone.

Stereotaxic surgery

Animals were anesthetized (90 mg/kg ketamine and 10 mg/kg xylazine, intraperitoneally) followed by analgesic (butorphanol) and antibiotic (gentamicin) administration. Rats received bilateral 1 µL microinjections (5 x 10⁶ tu/µL titer) into the IL (2.9 mm anterior to bregma, 0.6 mm lateral to midline, and 4.2 mm ventral from dura), as described previously (McKlveen et al., 2013; Myers et al., 2016, 2017), of either the vGluT1 siRNA virus or GFP control. All injections were carried out with a 25-gauge, 2-µL microsyringe (Hamilton, Reno, NV) using a microinjection unit (Kopf, Tujunga, CA) at a rate of 5 min/µL. To reduce tissue damage and allow diffusion, the needle was left in place for 5 minutes before and after injections. Animals recovered for 6 weeks before commencing experiments, corresponding to timeframes previously used for similar lentiviral systems (Grillo et al., 2007; Myers et al., 2017).
Telemetry

Four weeks after stereotaxic surgery, rats were implanted with radiotelemetry transmitters (PA-C40; Data Sciences International, St. Paul, MN) as previously described (Flak et al., 2011; Goodson et al., 2017). Briefly, animals were anesthetized with inhaled isoflurane anesthesia (1-5%). The descending aorta was exposed via an abdominal incision, allowing implantation of a catheter extending from the transmitter. The catheter was secured with tissue adhesive (Vetbond; 3M Animal Care Products, St. Paul, MN) and a cellulose patch. The transmitter body was then sutured to the abdominal musculature, followed by suturing of the abdominal musculature and closure of the skin with wound clips. Rats recovered for 2 weeks before wound clips were removed and experiments began.

Chronic variable stress

CVS was comprised of twice daily (AM and PM) repeated and unpredictable stressors, including exposure to a brightly-lit open field (1 m², 5 minutes), cold room (4°C, 1 hour), forced swim (23° to 27°C, 10 minutes), brightly-lit elevated platform (0.5 m, 5 minutes), shaker stress (100 rpm, 1 hour), and hypoxia (8% oxygen, 30 minutes). Additionally, overnight stressors were variably included, comprised of social crowding (6-8 rats/cage, 16 hours) and restricted housing (mouse cage, 16 hours). To prevent body weight differences between stress conditions, rats remaining unstressed in the home cage were food restricted in accordance with the reduced food intake induced by CVS (Flak et al., 2011, 2012). During the 2 weeks of CVS, unstressed animals received 80% of a food allotment prior to lights off and the other 20% after lights on to reduce the
potential for fasting (Flak et al., 2011, 2012). On day 15, all rats were exposed to a novel acute restraint to directly compare the effects of vGluT1 knockdown on cardiovascular responses to acute and chronic stress.

**Acute stress**

The morning after completion of CVS (approximately 16 hours after the last stress exposure), all animals were subjected to a novel 40-minute restraint. Stress response assessment was initiated between 08:00 and 09:00 hours. Animals were placed in well-ventilated Plexiglass restraint tubes with baseline pressure and HR measurements collected in the one-hour period preceding restraint. After restraint, rats were returned to their home cage with pressure and HR recorded for sixty minutes after restraint.

**Tissue collection**

After acute restraint, all animals were given an overdose of sodium pentobarbital and transcardially perfused with 0.9% saline followed by 4% phosphate-buffered paraformaldehyde. Brains were postfixed in paraformaldehyde for 24 hours and then stored in 30% sucrose at 4°C. Brains were subsequently sectioned (30 μm-thick coronal sections) and processed for GFP immunohistochemistry to determine microinjection spread.
**Immunohistochemistry**

For single immunolabeling of GFP, tissue sections were washed in 50 mM KPBS and incubated in blocking buffer (50 mM KPBS, 0.1% bovine serum albumin, and 0.2% TritonX-100) for 1 hour at room temperature. Sections were placed in rabbit anti-GFP primary antibody (1:1000 in blocking buffer; Invitrogen, La Jolla, CA) overnight at 4°C. Following incubation, sections were rinsed and placed into Alexa488-conjugated donkey anti-rabbit immunoglobulin G (IgG; 1:500 in blocking buffer; Jackson Immunoresearch, West Grove, PA) for 30 minutes. Sections were rinsed, mounted onto slides, and cover slipped. Dual fluorescent immunolabeling was performed as described previously (Myers et al., 2017), with GFP labeled in sequence with vGluT1. vGluT1 was visualized with rabbit anti-vGluT1 primary antibody (1:1000; Synaptic Systems, Goettingen, Germany) followed by Cy3-conjugated donkey anti-rabbit IgG (1:500; Jackson ImmunoResearch, West Grove, PA).

**Microscopy**

For visualization of GFP and vGluT1 co-localization, digital images were captured from a 1-in-12 series with a Zeiss Axio Imager Z2 microscope using optical sectioning (63x objective) to permit co-localization within a given z-plane (0.5-µm thickness). Co-localizations were defined as white fluorescence from overlap between labeled GFP terminals and magenta-colored vGluT1. For each figure, brightness and contrast were enhanced uniformly using Adobe Photoshop (CC 14.2).
Data analysis

Data are expressed as mean +/- standard error of the mean (SEM). All analyses were conducted using GraphPad Prism (version 7.04) for 2-way ANOVA, R Studio (version 3.4.2) for 3-way repeated measures ANOVA, or Dataquest A.R.T. (version 4.3) for telemetry analysis. Over the course of CVS, activity, HR, arterial pressures, and pulse pressure were sampled during the light phase from 06:00-08:00 for AM measures and during the dark phase from 19:00-21:00 for the PM period. During each 2-hour time period, samples were averaged into 10-minute bins for analysis. Beginning the day before CVS, circadian curves were generated for each parameter over the 15-day period. Data over time (both CVS and acute restraint) were analyzed by 3-way repeated measures analysis of variance (ANOVA); with viral treatment, stress, and time (repeated) as factors. When significant main effects were reported, ANOVA was followed by Tukey post-hoc test to identify specific group differences. Area under the curve (AUC) for the 15 days of CVS or 100 min of acute stress was calculated by summing the average values acquired between two time points multiplied by the time elapsed \[ \Sigma((\text{Time 1} + \text{Time 2})/2)\times\text{Time elapsed}\]. Statistical significance for cumulative measures was determined by 2-way ANOVA with treatment and stress as factors. ANOVA was followed by Tukey post-hoc tests in the case of significant main effects. Values more than 2 standard deviations from the mean were removed as outliers. Statistical significance was reported as (p < 0.05) for all tests.
Experiment 2

Design

Experiment 2 employed a similar design as experiment 1. Male rats (n = 7/group) received IL injections of lentiviral-packaged vGluT1 siRNA or GFP and experienced CVS or remained unstressed. On day 15, thoracic aorta was collected to examine vasoreactivity and histology. Additionally, hearts and brains were collected for histological analyses.

Chronic variable stress

For experiment 2, CVS was staggered based on the throughput of vascular function experiments. Seven groups (n = 4 animals, one of each treatment) began the CVS paradigm one day apart so that each group of 4 would have tissue collection on successive days. All rats undergoing CVS in experiment 2 received the same stressors in the same sequence. The CVS paradigm was similar to experiment 1, except for the substitution of restraint (30 min) for crowding.

Tissue collection

The morning of day 15, approximately 16 hours after the last stress exposure, all animals were rapidly anesthetized with isoflurane and decapitated. Thoracic aorta was collected by dissecting 4 mm of aortic tissue proximal to the diaphragm for vascular function analysis. Additional aortic tissue was collected proximal to the initial sample and post-fixed in paraformaldehyde for histological analysis. Hearts were also collected and post-fixed in paraformaldehyde for histological analysis. Brains were post-fixed and
subsequently sectioned and processed for GFP immunohistochemistry to determine microinjection spread as described for experiment 1.

**Vascular function**

Aortic tissue samples were processed for wire myographic vasoreactivity analyses as previously described (Basford *et al.*, 2013). Briefly, vessels were kept in warm, oxygenated Krebs’s solution with connective tissue and adipose removed under a dissecting microscope. Aortic rings were then placed on wires in organ baths (Radnoti, Covina, CA), equilibrated, and brought to tension. Vasoconstriction was assessed in response to increasing concentrations of the endothelial-dependent potassium chloride (KCl; 0 to 50 mM) and endothelial-independent phenylephrine (1x10^{-6} to 1x10^{-2} mM). Vessels were then set to 80% of maximum phenylephrine-induced constriction and relaxation was determined in response to endothelial-dependent acetylcholine (1x10^{-6} to 1x10^{-2} mM) and endothelial-independent sodium nitroprusside (SNP; 1x10^{-7} to 3x10^{-3} mM). At the end of the experiment, all vessels were weighed and measured (length, circumference, and area) to verify equal dimensions across all groups.

**Histology and microscopy**

Cardiac and aortic tissue was processed by the Cincinnati Children’s Hospital Medical Center Research Pathology Core. Briefly, aortas were paraffin-embedded, sectioned (5 µm), and stained. Verhoeff-Van Gieson (VVG) was used to quantify elastin in the tunica media and Masson’s Trichrome to quantify collagen in the tunica adventitia.
as previously described (Basford et al., 2013; Goodson et al., 2017). Paraffin-embedded hearts were oriented for four-chamber view, sectioned (5 \( \mu \)m), and stained with Masson’s Trichrome to visualize collagen and wheat germ agglutinin (WGA) conjugated to Alexa488 to visualize myocyte cell membranes as previously described (Basford et al., 2013; Gupta et al., 2016). Vascular and heart tissue were imaged with a Zeiss AxioObserver microscope using a color camera and 10x objective.

**Data analysis**

Data are expressed as mean ± SEM. All analyses were conducted using GraphPad Prism (version 7.04) for 2-way ANOVA, R Studio (version 3.4.2) for 3-way repeated measures ANOVA, or FIJI (version 1.51N) for histological quantification. Vasoreactivity data were analyzed by 3-way repeated measures ANOVA; with viral treatment, stress, and drug concentration (repeated) as factors. When significant main effects were reported, ANOVA was followed by Tukey post-hoc test to identify specific group differences. FIJI (version 1.51N) was used to quantify lumen and tunica media dimensions in VVG-stained aorta, as well as adventitia dimensions in Masson’s Trichrome-stained tissue. FIJI was also used to quantify collagen density in hearts stained with Masson’s Trichrome. For each animal, 6 sections of aorta or heart were quantified and averaged. To determine myocyte surface area, FIJI was used to binarize myocyte images. This technique produced dark cytoplasm and bright membranes in cardiomyocytes (Bensley et al., 2016). The dark cytoplasm was used to calculate surface areas of the myocytes in the apex and lateral wall of the left ventricle. For each animal, approximately 100 cells were counted from 4 heart sections and averaged.
Statistical significance for histological measures was determined by 2-way ANOVA with treatment and stress as factors. ANOVA was followed by Tukey post-hoc tests in the case of significant main effects. Values more than 2 standard deviations from the mean were removed as outliers. Statistical significance was reported as (p < 0.05) for all tests.

Results

vGluT1 knockdown

Injections of a lentiviral-packaged construct expressing vGluT1 siRNA were targeted to the IL (Fig. 1a, 1b). Viral injections were largely limited to the deep layers of IL, with minimal spread to PL. We have previously shown that this approach reduces vGluT1 mRNA specifically in the IL, as well as vGluT1 protein co-localization with GFP-labeled terminals (Myers et al., 2017). In the current study, tissue from rats injected with a GFP control construct exhibited substantial co-localization with vGluT1 protein on cortico-cortical axonal processes (Fig. 1c). IL projections transfected with the vGluT1 siRNA construct had reduced co-localization with vGluT1 protein (Fig. 1d).

Body weight and food intake

Chronic stress reduces food intake and body weight gain, leading to significant differences in body composition compared to control animals (Solomon et al., 2010; Flak et al., 2011; Smith et al., 2017; Myers et al., 2017). As this may confound results related to HR and blood pressure reactivity (Flak et al., 2011), animals in the No CVS groups for both experiments 1 and 2 received mild food restriction to match body weight with CVS rats (Table 1). In both experiments, there were no significant differences in
body weight between groups. However, food restriction in experiment 1 led to food consumption that was significantly greater than the CVS groups [F(1,28) = 31.24, p < 0.0001]. In experiment 2, food restriction significantly decreased food intake compared to CVS groups [F(1,24) = 33.25, p < 0.0001].

Experiment 1

Circadian behavioral activity

Home cage radiotelemetry data were analyzed for 15 days beginning the day before CVS. Throughout CVS, 3-way repeated-measures ANOVA of circadian activity (n = 7-8/group) revealed a main effect of time [F(29, 930) = 21.38, p < 0.0001] and interactions of stress x treatment [F(1, 930) = 4.67, p = 0.031] and stress x time [F(29, 930) = 4.34, p < 0.0001]. Post-hoc analysis indicated that, during the dark period of CVS day 2, rats that were subjected to an overnight crowding stressor exhibited more activity than No CVS controls (Fig. 2a). The CVS GFP rats had elevated activity compared to No CVS GFP controls (p = 0.014); furthermore, CVS siRNA animals were more active than CVS GFP rats (p = 0.007). In addition to overnight social crowding on day 2, rats experienced overnight housing in mouse cages days 6 and 10. Throughout the dark periods of days 6-9, CVS siRNA rats had decreased activity compared to No CVS animals (p ≤ 0.047). Additionally, the CVS GFP rats were less active on the dark periods of days 9 (p = 0.008) and 14 (p = 0.003) compared to No CVS GFP. Also, on the dark period of Day 14, the CVS siRNA group was more active than CVS GFP (p = 0.003). Additional analysis was carried out on cumulative activity counts (No CVS GFP: 801.88 ± 1.04, No CVS siRNA: 819.29 ± 8.08, CVS GFP: 683.66 ±1.38, CVS siRNA: 703.44 ± 1.25).
860.81 ± 4.46). According to 2-way ANOVA, there were significant effects of treatment 
[F(1,27) = 338.60, p < 0.0001], stress [F(1,27) = 60.38, p < 0.0001], and a treatment x 
stress interaction [F(1,27) = 261.9, p < 0.0001]. Tukey post-hoc test indicated that total 
activity over 15 days was lower in CVS GFP rats compared to No CVS GFP (p < 
0.0001). In contrast, cumulative activity was increased in CVS siRNA rats compared to 
all other groups (p < 0.0001).

Circadian heart rate

Home cage HR (n = 8/group) was analyzed by 3-way repeated-measures 
ANOVA revealing a main effect of time, [F(29, 960) = 72.64, p < 0.0001] accompanied 
by a stress x treatment interaction [F(1, 960) = 28.71, p < 0.0001]. Early in CVS (Fig. 
2b), the CVS siRNA group had elevated dark phase HR compared to the CVS GFP 
group (Day 0, p = 0.024), as well as the No siRNA group during overnight social 
crowding (Day 2, p = 0.024). Also, on the first light period of CVS, the CVS GFP HR 
was decreased compared to No CVS GFP (p = 0.028).

Circadian and cumulative arterial pressures

In order to examine the effects of chronic stress and decreased IL output on 
long-term blood pressure regulation, home cage arterial pressures (n = 6-8/group) were 
continuously monitored. For mean arterial pressure (MAP), 3-way repeated-measures 
ANOVA found main effects of stress [F(1, 900) = 15.33, p < 0.0001] and time [F(29, 
900) = 13.91, p < 0.0001], as well as an interaction of stress x treatment [F(1, 900) = 
14.41, p < 0.0001]. In terms of time-specific effects, there was an increase in MAP in the 
CVS siRNA group compared to No CVS siRNA (p = 0.022, Fig. 3a) during the light
period of CVS day 1. While few time point-specific circadian effects were identified, AUC analysis found that the CVS siRNA group experienced greater cumulative MAP (p < 0.0001, Fig. 3b) and systolic arterial pressure (SAP, p < 0.0001, Fig. 3c) than all other groups. Diastolic arterial pressure (DAP, p < 0.0001, Fig. 3d) was also elevated in the CVS siRNA animals compared to No CVS siRNA and CVS GFP. Between non-stressed rats, the siRNA treatment led to lower cumulative pressures (MAP, SAP, and DAP, p < 0.0001) relative to GFP.

Circadian and cumulative pulse pressure

Pulse pressure is a function of vascular stiffness and predicts heart disease independent of MAP (Franklin et al., 1999; Glasser et al., 2014). Three-way repeated-measures ANOVA identified a main effect of stress [F (1, 900) = 34.35, p < 0.0001] and an interaction of stress x treatment [F(29, 900 = 2.59, p < 0.0001] for circadian pulse pressure. Over the course of CVS, the only time-specific difference in circadian pulse pressure occurred during the dark phase of day 0 (Fig. 4a) where the CVS GFP pulse pressure was greater than No CVS GFP (p = 0.01, n = 6-8/group). Cumulative pulse pressure from AUC analysis was increased in both CVS groups relative to No CVS (p < 0.0001, Fig 4b). Additionally, CVS siRNA cumulative pulse pressure was greater than all groups, including CVS GFP (p < 0.0001).

Acute stress reactivity

In order to study the role of the IL in acute stress reactivity, MAP and HR reactivity were monitored during restraint (n = 8/group). During acute stress, 3-way
repeated-measures ANOVA of HR reactivity found a main effect of time \[ F(1, 84) = 53.99, p < 0.0001 \] (Fig. 5a). Compared to No CVS GFP, vGluT1 siRNA elevated HR during restraint minutes 15-25 \( (p \leq 0.040) \). Both CVS GFP and CVS siRNA rats had elevated HR compared to No CVS GFP from 10-25 minutes of restraint \( (p \leq 0.006) \). Further, CVS siRNA rats had elevated HR during restraint at 35 and 40 minutes \( (p \leq 0.008) \). While recovering from stress in the home cage, HR remained elevated in the CVS siRNA group relative to No CVS GFP at minutes 45 and 95 \( (p \leq 0.010) \).

Additionally, CVS GFP and No CVS siRNA had elevated HR on minutes 45 \( (p = 0.0003) \) and 95 \( (p = 0.031) \), respectively. Cumulative HR reactivity from AUC analysis of acute stress responses revealed that both No CVS siRNA and CVS GFP groups had elevated HR responses to acute restraint \( (p < 0.0001, \text{Fig. 5b}) \). Moreover, the CVS siRNA group experienced greater cumulative HR than all other groups \( (p = 0.0006) \).

Analysis of stress-evoked MAP by 3-way repeated-measures ANOVA identified a main effect of time \[ F(1, 84) = 99.91, p < 0.0001 \] (Fig. 5c). The CVS GFP group had greater MAP reactivity compared to No CVS GFP on minutes 5-15 of restraint \( (p \leq 0.030) \). The CVS siRNA animals had greater MAP than No CVS GFP at 15, 20, and 40 minutes \( (p \leq 0.05) \). During recovery, CVS GFP MAP remained elevated at 50, 75, and 80 minutes \( (p \leq 0.04) \); furthermore, CVS siRNA MAP was higher at minutes 75-90 \( (p \leq 0.045) \). AUC analysis by 2-way ANOVA found siRNA treatment increased cumulative MAP \( (p < 0.0001, \text{Fig. 5d}) \). CVS also increased MAP AUC as both CVS groups were higher than respective No CVS controls \( (p < 0.0001) \).
**Experiment 2**

**Injection placement**

Similar to experiment 1, injections of lentiviral-packaged constructs were targeted to the IL with minimal spread to PL (Fig. 6). Although, injections from experiment 2 had greater spread into superficial layers of the IL. Additionally, some injections spread into the striatum caudally but this region does not exhibit vGluT1 expression (Ziegler et al., 2002).

**Vascular function**

In order to assess the vascular consequences of prolonged stress, arterial function was monitored in response to endothelial-dependent and-independent agents ex vivo. In terms of endothelial-independent vasoconstriction, 3-way repeated-measures ANOVA identified a main effect of KCl concentration [$F(1,36) = 669.73, p < 0.0001$] and an interaction of drug concentration x viral treatment [$F(1,36) = 9.55, p = 0.004$] (Fig. 7a). As determined by post-hoc analyses, aortas of CVS siRNA animals constricted less than No CVS GFP at KCl concentrations above of 25 mM ($p \leq 0.0006$). CVS siRNA also showed impaired constriction compared to CVS GFP at concentrations above 30 mM ($p \leq 0.006$). Compared to No CVS siRNA, CVS siRNA vasoreactivity was decreased at concentrations of 30 and 50 mM ($p \leq 0.042$). Within the No CVS groups, siRNA treatment decreased constriction at 40 mM ($p = 0.019$). Endothelium-dependent vasoconstriction in response to phenylephrine showed a main effect of concentration [$F(1,40) = 31.62, p < 0.0001$] (Fig. 7b). Either siRNA or CVS alone impaired vasoreactivity at drug concentrations above 1 $\mu$M ($p < 0.05$). However, CVS siRNA
animals had impaired vasoconstriction compared to all other groups at phenylephrine concentrations above 0.1 µM (p < 0.05).

Endothelial-independent vasorelaxation to acetylcholine showed a main effect of drug concentration [$F(1,40) = 15.89, p = 0.0003$] by 3-way repeated-measures ANOVA (Fig. 7c). Post-test found that CVS siRNA vasorelaxation was decreased compared to all groups at acetylcholine concentrations above 1 µM (p = 0.048). Endothelial-dependent vasorelaxation to SNP, as analyzed by 3-way repeated-measures ANOVA, showed a main effect of SNP concentration [$F(1,44) = 12.65, p = 0.0009$] (Fig. 7d). CVS siRNA tissue had impaired vasorelaxation compared to both No CVS groups at concentrations above of 0.03 µM (p = 0.026). Additionally, CVS alone impaired vasorelaxation in GFP rats at SNP concentrations above 0.3 µM (p < 0.010).

Vascular and cardiac histology

Histological analysis was carried out to investigate the effects of chronic stress and vGluT1 knockdown on markers of vascular and cardiac pathology (Table 2). In GFP-injected rats, CVS increased aortic tunica media thickness [$F(1,24) = 18.55, p = 0.0002$] and media:lumen area [$F(1,24) = 10.45, p < 0.004$]. Furthermore, CVS increased adventitial fibrosis in terms of increased collagen density [$F(1,24) = 4.944, p = 0.036$]. Chronic stress also affected the myocardium by increasing heart weight [$F(1,24) = 7.028, p = 0.014$] and myocyte surface area [$F(1,23) = 5.084, p = 0.034$]. In animals with reduced vGluT1, CVS had greater effects on aortic remodeling. CVS siRNA rats had decreased luminal circumference [$F(1,24) = 8.217, p = 0.022$] and area [$F(1,24) = 8.142, p = 0.026$], increased media thickness [$F(1,24) = 18.55, p = 0.026$] and
media:lumen area \[ F(1,24) = 10.45, p < 0.0004 \], increased collagen density \[ F(1,24) = 4.944, p = 0.036 \] and decreased adventitial thickness \[ F(1,24) = 6.517, p = 0.031 \] (Fig. 8a-d). Collectively, these results indicate that CVS interacts with decreased IL function to promote fibrosis and inward remodeling of vascular muscle leading to restricted luminal area, potentially accounting for impaired vasoreactivity and arterial stiffness. CVS siRNA rats also exhibited cardiac hypertrophy as these animals had increased heart weight \[ F(1,24) = 7.028, p = 0.014 \], heart weight relative to body weight \[ F(1,24) = 17.09, p = 0.015 \], and increased myocyte surface area \[ F(1,23) = 5.084, p = 0.034 \] (Fig. 8e,f), without affecting myocardial collagen deposition.

**Discussion**

In the current study, we utilized viral-mediated gene transfer to decrease IL glutamatergic output while simultaneously monitoring hemodynamic, vascular, and cardiac responses to chronic stress. We found that IL vGluT1 knockdown increased heart rate and arterial pressure reactivity to acute stress. Additionally, IL hypofunction during CVS increased chronic home cage arterial pressure. These changes were accompanied by both endothelial-independent and -dependent arterial dysfunction. Histological analysis revealed that animals experiencing CVS with decreased IL output also had inward vascular remodeling, fibrosis, and cardiac hypertrophy. Collectively, these results indicate that IL projection neurons are critical for reducing acute cardiovascular stress reactivity, long-term arterial pressure, and vascular endothelial dysfunction. Furthermore, they identify a neurochemical mechanism linking stress appraisal and emotion with chronic stress-induced autonomic dysfunction.
Epidemiological evidence indicates that prolonged stress is a major risk factor for cardiovascular illness and mortality (Yusuf et al., 2004; Steptoe & Kivimäki, 2012). Additionally, numerous clinical studies point to enhanced stress reactivity as a marker of future cardiovascular pathology (Chida & Steptoe, 2010; Vogelzangs et al., 2010). Rodent studies employing repeated-stress models of depression (chronic variable stress, chronic mild stress, chronic social defeat, etc.) have found alterations in baroreflex function, decreased heart rate variability, and ventricular arrhythmias (Grippo et al., 2002; Costoli, 2004; Grippo & Johnson, 2009; Wood et al., 2012; Carnevali et al., 2013; Wood, 2014; Duarte et al., 2015; Crestani, 2016). Given the strong association between prolonged stress/emotional disorders and cardiovascular disease (Carney et al., 2001; Barton et al., 2007; Hausberg et al., 2007), it is important to identify the specific neural processes of stress appraisal and mood that impact cardiovascular function. Prolonged exposure to stress mediators related to autonomic imbalance generates risk for cardiovascular pathology (Johnson & Grippo, 2006.; Thayer et al., 2012; Wulsin et al., 2015). Stress-associated molecules such as corticosteroids, corticotropin-releasing hormone, and neuropeptide Y, among others, have been shown to affect cardiovascular function in animal models (Costoli et al., 2005; Wood et al., 2012; Oakley et al., 2019). However, the neural circuits that integrate cognitive appraisal and autonomic balance remain to be determined.

The contribution of the current study relates to the site-specific genetic approach that reduces IL vGluT1 expression long-term (Myers et al., 2017). Given the decreased output from a critical cognitive/emotional region, we monitored both stress-induced and resting parameters of heart rate and arterial pressure in otherwise unstressed rats, as
well as rats experiencing the cumulative burden of chronic stress exposure. This approach was followed by ex vivo analysis of arterial function and histological investigation of vascular and myocardial structure. These studies found several siRNA effects in animals that did not experience CVS (Table 3). The knockdown increased acute HR and MAP reactivity to restraint and impaired endothelial-independent vasoconstriction, suggesting hypo-function of IL primes for enhanced cardiovascular reactivity and impaired vasomotor function. Paradoxically, these animals exhibited decreased resting home cage circadian arterial pressure. This effect may relate to the important role of IL molecular clocks in coordinating circadian physiological rhythms (Woodruff et al., 2016). In GFP-treated rats, CVS dampened behavioral activity in the dark cycle, decreasing circadian rhythms of activity and potentially accounting for effects of CVS to decrease circadian HR. Interestingly, CVS did not alter chronic home cage MAP; however, CVS-exposed animals exhibited enhanced tachycardic and pressor responses to restraint. This was accompanied by impaired endothelial-independent vasorelaxation and constriction. Furthermore, CVS increased vascular smooth muscle thickness and fibrosis, as well as cardiomyocyte surface area.

The effects of vGluT1 knockdown and CVS interacted to generate a phenotype of enhanced cardiovascular risk. All measures of HR and arterial pressure, both acute stress-induced and chronic home cage, were elevated. This group also had impairment of both endothelial-dependent and –independent vascular dilation and constriction. The overall risk profile was further evident by inward remodeling of the vasculature due to hypertrophy and fibrosis that reduced luminal area, indicative of vascular stiffness. Additionally, these rats exhibited myocardial hypertrophy including increased myocyte
size, heart weight, and body weight-corrected heart weight. Taken together, these results suggest that conditions associated with decreased ventral mPFC activity, including depression and anxiety, may enhance vulnerability to the effects of prolonged stress on cardiovascular health.

The specific circuit mechanisms downstream of IL glutamate release that account for the current findings remain to be determined. Although the IL does not directly innervate either pre-ganglionic autonomic neurons or neurosecretory cells of the hypothalamus (Saper et al., 1976; Vertes, 2004), the region has widespread projections throughout the forebrain and brainstem (Vertes, 2004; Wood et al., 2019). Inputs to the posterior hypothalamus target local inhibitory cells (Myers et al., 2016), providing a potential pathway to inhibit stress reactivity (Lisa et al., 1989). There are also IL projections to brainstem cardioregulatory centers including the nucleus of the solitary tract and the rostral and caudal regions of the ventrolateral medulla (Gabbott et al., 2005; Myers, 2017). Ultimately, circuit-specific analyses may isolate the precise cell populations within IL glutamate neurons that reduce autonomic imbalance after chronic stress. Additionally, there is a complex network of interneurons in the IL that mediates the overall activity of glutamatergic projection neurons (McKlveen et al., 2019); accordingly, these local circuits would be expected to play a role in autonomic reactivity.

While these studies identified a novel fronto-cortical node for preventing the deleterious cardiovascular effects of chronic stress, there are limitations worth discussing. First, the current studies were limited to males. As depression-cardiovascular co-morbidity has twice the prevalence in females (Möller-Leimkühler, 2007; Tobet et al., 2013), it is important to consider sex-specific regulation.
a recent study investigating vascular function in female rodents after chronic stress actually found that ovarian hormones protect against stress-induced arterial dysfunction (Brooks et al., 2018). Comparing the neural basis of pathological responses in males and females would likely yield a better understanding of the disproportionate female impact of mood disorder-cardiovascular co-morbidity. Another consideration is that the current vasoreactivity experiments were carried out in thoracic aorta. Although the results identified impaired function and indicators of stiffness, the aorta is a conductive artery and future experiments with resistance arterioles might yield differing results. Indeed, these studies could show greater effects as resistance arteries receive more sympathetic innervation (Hao et al., 2005; Brown et al., 2018). Furthermore, investigating vascular resistance could yield data with significant relevance for stress-related hypertension.

In conclusion, the current findings highlight IL glutamatergic neurons as a node of integration that links stress appraisal with hemodynamic reactivity, long-term arterial pressure control, and vascular endothelial function. These results also indicate that, in the context of chronic stress, cortical cells mediating cognition and behavior can impact the structure and function of the heart and vasculature. Future research investigating the mechanisms that regulate IL projection neuron activity and the downstream post-synaptic events activated by IL glutamate release may yield insight into novel targets to prevent or reduce the burden of cardiovascular disease.
Acknowledgments

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Disclosures

J. M. McKlveen contributed to this article in her personal capacity. The views expressed herein are those of the authors and do not necessarily represent the views of the National Institutes of Health, National Center for Complimentary and Integrative Health, or the United States Government. All authors report no biomedical financial interests or potential conflicts of interest.
**Figure Legends**

**Figure 1.** The spread of individual lentiviral injections was traced on photomicrographs and overlaid onto atlas templates from Swanson (Swanson, 2004) to depict the localization of vGluT1 knockdown in experiment 1 (a). Lentiviral injections targeted to the IL with minimal spread to the PL (b). White arrows indicate dorsal and ventral boundaries of the IL. Immunolabeling of GFP (green) and vGluT1 (magenta) indicated a high-degree of co-localization (white arrows) on IL projections in GFP controls (c). Knockdown of vGluT1 with siRNA treatment decreased vGluT1 co-localization with GFP on IL projections (d). Scale bars: (b) 100 µm, (c,d) 10 µm. Numbers indicate distance rostral to bregma in millimeters. AC: anterior cingulate, PL: prelimbic cortex, IL: infralimbic cortex, fa: anterior forceps of the corpus callosum.

**Table 1.** Body weight of animals at the beginning and end of CVS for experiments 1 and 2. In both experiments, CVS rats had *ad libitum* access to chow while No CVS animals received mild food restriction to prevent significant differences in body weight between chronically stressed animals and controls.

**Figure 2.** Chronically stressed rats had disrupted circadian behavioral rhythms evidenced by interactions of CVS and siRNA, as well as CVS and time (n = 7-8/group) (a). Circadian heart rate also exhibited a CVS x siRNA interaction (n = 8/group) leading to heart rate disruptions early in CVS (b). *p < 0.05 CVS GFP vs. No CVS GFP, ^p < 0.05 CVS siRNA vs. No CVS GFP, †p < 0.05 CVS siRNA vs. CVS GFP, &p < 0.05 CVS siRNA vs. No CVS siRNA.
**Figure 3.** Chronic stress and siRNA treatment (n = 8/group) interacted leading to altered circadian arterial pressure (a). Analysis of cumulative arterial pressure indicated that CVS, only in siRNA-treated rats, increased chronic MAP (b), SAP (c), and DAP (d). *p < 0.05 No CVS siRNA vs. No CVS GFP, ^p < 0.05 CVS siRNA vs. No CVS GFP, †p < 0.05 CVS siRNA vs. CVS GFP, &p < 0.05 CVS siRNA vs. No CVS siRNA.

**Figure 4.** Chronic stress and siRNA treatment (n = 6-8/group) interacted to affect circadian pulse pressure (a). Cumulative pulse pressure was increased in CVS exposed rats with CVS siRNA rats experiencing the greatest chronic pulse pressure (b). #p < 0.05 CVS GFP vs. No CVS GFP, ^p < 0.05 CVS siRNA vs. No CVS GFP, †p < 0.05 CVS siRNA vs. CVS GFP, &p < 0.05 CVS siRNA vs. No CVS siRNA.

**Figure 5.** In response to acute restraint, both siRNA and CVS (n = 8/group) increased heart rate reactivity and impaired recovery (a). Cumulative acute heart rate responses were also elevated by siRNA and CVS but CVS siRNA rats had the greatest overall heart rate response (b). Chronically stressed rats, both GFP and siRNA treated, had increased MAP reactivity and impaired recovery (c). Analysis of cumulative restraint-induced pressor responses indicated effects of both siRNA and CVS (d). *p < 0.05 No CVS siRNA vs. No CVS GFP, #p < 0.05 CVS GFP vs. No CVS GFP, ^p < 0.05 CVS siRNA vs. No CVS GFP, †p < 0.05 CVS siRNA vs. CVS GFP, &p < 0.05 CVS siRNA vs. No CVS siRNA.
**Figure 6.** The spread of individual lentiviral injections was traced on photomicrographs and overlaid onto atlas templates from Swanson (Swanson, 2004) to depict the localization of vGluT1 knockdown in experiment 2. White arrows indicate dorsal and ventral boundaries of the IL. Numbers indicate distance rostral to bregma in millimeters. AC: anterior cingulate, PL: prelimbic cortex, IL: infralimbic cortex.

**Figure 7.** Aortic tissue from CVS siRNA animals (n = 7/group) had impaired endothelial-dependent vasoconstriction (a). Both CVS and siRNA impaired endothelial-independent vasoreactivity but the CVS siRNA group had the greatest impairment (b). Endothelial-dependent vasorelaxation was impaired only in the CVS siRNA group (c), while endothelial-independent relaxation was impaired in CVS GFP and CVS siRNA tissue (d). *p < 0.05 No CVS siRNA vs. No CVS GFP, †p < 0.05 CVS GFP vs. No CVS GFP, ‡p < 0.05 CVS siRNA vs. No CVS GFP, §p < 0.05 CVS siRNA vs. No CVS siRNA.

**Figure 8.** Verhoeff-van Gieson stain was used to visualize elastin (dark brown) in aortic tissue of No CVS GFP (a) and CVS siRNA (b) rats. Greater thickness of the tunica media is indicated by white arrows. Masson’s Trichrome was used to stain collagen (blue) in aortic tissue of No CVS GFP (c) and CVS siRNA (d) animals. White arrows indicate increased collagen density in the tunica adventitia. Wheat germ agglutinin conjugated to Alexa 488 (green) was used to visualize cardiomyocyte membranes (e). Binarized myocyte images (f) were used to quantify myocyte surface area. Scale bars: (a-d) 50 µm, (e) 10 µm. * denotes the lumen.
Table 2. Histological quantification of vascular and myocardial structure. Dimensions of the lumen and media were quantified from elastin staining. Adventitial size and fibrosis were determined from collagen staining. Myocyte surface area was measured with membrane labeling and cardiac fibrosis was queried via collagen staining. There were no effects of siRNA alone. CVS increased media thickness, adventitial collagen, and myocyte size. In contrast, CVS siRNA tissue exhibited alterations in all structural endpoints assessed except myocardial fibrosis.

Table 3. Integrative summary of all data reported in terms of siRNA effects, chronic stress effects, and the combination of siRNA and CVS.
References


Stimulated release of tissue plasminogen activator from artery wall sympathetic nerves: Implications for stress-associated wall damage. *Stress* 8, 141–149.


Myers B, McKlveen JM, Morano R, Ulrich-Lai YM, Solomon MB, Wilson SP & Herman


Fig 3

A  Circadian Mean Arterial Pressure

B  Cumulative Mean Arterial Pressure

C  Cumulative Systolic Arterial Pressure

D  Cumulative Diastolic Arterial Pressure
Fig 4
<table>
<thead>
<tr>
<th>Group</th>
<th>Body Weight CVS Day 1 (g)</th>
<th>Body Weight CVS Day 13 (g)</th>
<th>Daily Food Intake (g)</th>
</tr>
</thead>
<tbody>
<tr>
<td>No CVS GFP</td>
<td>389.78 ± 6.21</td>
<td>405.06 ± 6.14</td>
<td>21.04 ± 0.36</td>
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<tr>
<td>No CVS siRNA</td>
<td>389.84 ± 12.23</td>
<td>405.85 ± 12.13</td>
<td>21.68 ± 0.53</td>
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<tr>
<td>CVS GFP</td>
<td>402.03 ± 7.45</td>
<td>404.19 ± 8.57</td>
<td>18.82 ± 0.30*#</td>
</tr>
<tr>
<td>CVS siRNA</td>
<td>417.16 ± 6.79</td>
<td>417.64 ± 6.25</td>
<td>18.89 ± 0.35*&amp;</td>
</tr>
</tbody>
</table>

**Experiment 1 n = 8/group**

<table>
<thead>
<tr>
<th>Group</th>
<th>Body Weight CVS Day 1 (g)</th>
<th>Body Weight CVS Day 13 (g)</th>
<th>Daily Food Intake (g)</th>
</tr>
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<tbody>
<tr>
<td>No CVS GFP</td>
<td>372.83 ± 7.72</td>
<td>363.80 ± 7.15</td>
<td>15.08 ± 0.45</td>
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<tr>
<td>No CVS siRNA</td>
<td>379.19 ± 5.03</td>
<td>368.00 ± 4.73</td>
<td>15.35 ± 0.29</td>
</tr>
<tr>
<td>CVS GFP</td>
<td>381.44 ± 9.07</td>
<td>378.10 ± 9.33</td>
<td>17.44 ± 0.53#</td>
</tr>
<tr>
<td>CVS siRNA</td>
<td>384.69 ± 5.10</td>
<td>379.49 ± 3.64</td>
<td>17.78 ± 0.35*&amp;</td>
</tr>
</tbody>
</table>

* p < 0.05 CVS GFP vs. No CVS GFP, ^p < 0.05 CVS siRNA vs. No CVS GFP, Ȗ p < 0.05 CVS siRNA vs. No CVS siRNA

**Table 1. Body weight and food intake throughout chronic variable stress**
<table>
<thead>
<tr>
<th>Table 2. Vascular and cardiac histological analysis</th>
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<tbody>
<tr>
<td>n = 7/group</td>
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<td></td>
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<tr>
<td>Luminal Circumference (mm)</td>
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<tr>
<td>No CVS GFP</td>
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<tr>
<td>No CVS siRNA</td>
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<tr>
<td>CVS GFP</td>
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<tr>
<td>CVS siRNA</td>
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<tr>
<td>Luminal Area (mm^2)</td>
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<tr>
<td>No CVS GFP</td>
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<tr>
<td>No CVS siRNA</td>
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<tr>
<td>CVS GFP</td>
</tr>
<tr>
<td>CVS siRNA</td>
</tr>
<tr>
<td>Media Thickness (mm)</td>
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<tr>
<td>No CVS GFP</td>
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<tr>
<td>No CVS siRNA</td>
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<tr>
<td>CVS GFP</td>
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<tr>
<td>CVS siRNA</td>
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<tr>
<td>Media:Lumen Area</td>
</tr>
<tr>
<td>No CVS GFP</td>
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<tr>
<td>No CVS siRNA</td>
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<tr>
<td>CVS GFP</td>
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<tr>
<td>CVS siRNA</td>
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<tr>
<td>Adventitia Collagen (% area)</td>
</tr>
<tr>
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<tr>
<td>No CVS siRNA</td>
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<tr>
<td>CVS GFP</td>
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<tr>
<td>CVS siRNA</td>
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<tr>
<td>Adventitia Thickness</td>
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<tr>
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<td>No CVS siRNA</td>
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<tr>
<td>CVS GFP</td>
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<tr>
<td>CVS siRNA</td>
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<tr>
<td>Heart Weight (g)</td>
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<tr>
<td>No CVS siRNA</td>
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<tr>
<td>CVS GFP</td>
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<tr>
<td>CVS siRNA</td>
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<tr>
<td>Heart Weight/Body Weight (x100)</td>
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<tr>
<td>No CVS siRNA</td>
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<tr>
<td>CVS GFP</td>
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<tr>
<td>CVS siRNA</td>
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<td>Myocyte Surface Area (mm^2)</td>
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<td>No CVS siRNA</td>
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<tr>
<td>CVS GFP</td>
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<tr>
<td>CVS siRNA</td>
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<tr>
<td>Myocardial Collagen (% area)</td>
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<td>No CVS siRNA</td>
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<td>CVS GFP</td>
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<td>CVS siRNA</td>
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*p < 0.05 CVS GFP vs. No CVS GFP, ^p < 0.05 CVS siRNA vs. No CVS GFP, ^a p < 0.05 CVS siRNA vs. No CVS siRNA
<table>
<thead>
<tr>
<th>Table 3. Summary: siRNA and/or CVS effects from experiments 1 and 2.</th>
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<tr>
<td><strong>Activity (15 days)</strong></td>
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<tr>
<td><strong>HR (15 days)</strong></td>
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<td><strong>MAP (15 days)</strong></td>
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<td><strong>SAP (15 days)</strong></td>
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<td><strong>DAP (15 days)</strong></td>
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<td><strong>Pulse Pressure (15 days)</strong></td>
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<tr>
<td><strong>HR (acute stress)</strong></td>
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<td><strong>MAP (acute stress)</strong></td>
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<td><strong>Vasoconstriction (ED)</strong></td>
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<td><strong>Vasoconstriction (EI)</strong></td>
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<td><strong>Vasorelaxation (ED)</strong></td>
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<td><strong>Vasorelaxation (EI)</strong></td>
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<td><strong>Media Thickness</strong></td>
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<td><strong>Heart Weight</strong></td>
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<tr>
<td><strong>Heart Weight/Body Weight</strong></td>
</tr>
<tr>
<td><strong>Myocardial Collagen</strong></td>
</tr>
<tr>
<td><strong>Myocyte Surface Area</strong></td>
</tr>
</tbody>
</table>

Arrows indicate significant siRNA and/or CVS effects. HR: heart rate, MAP: mean arterial pressure, SAP: systolic arterial pressure, DAP: diastolic arterial pressure, ED: endothelial-dependent, EI: endothelial-independent.

*p < 0.05 vs No CVS GFP, *p < 0.05 vs No CVS siRNA, **p < 0.05 vs CVS GFP